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Date of Application and filing Complete Specification: 5-Aug., 1966 No. 48996/67.

(Patent of Addition to No. 1,112,033 dated 10 March, 1965).

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Phosphoric Acid Recovery.

Page 1, Heading Inventors: - for "AVRAGHAM MATITIAJU BANIEL" read "AVRAHAM MATITIAHU BANIEL"

THE PATENT OFFICE

containing phosphoric acid. The invention ture (that is, the solubility of phosphoric may be used for extracting phosphoric acid acid in the solvent is greater the lower the from a "technical-grade phosphoric_acid", i.e. an aqueous phosphoric acid, usually manufactured by the decomposition of rock phosphate::with::sulphuric::acid,_having:_an-H₂PO,_concentration=of=not=less=than=35% by_weight: and_possibly_up_to=about=90%= by weight, and containing the usual impuri-

ties of such phosphoric acid.
Our co-pending Application No. 10235/65
(Serial=No. 1,112,033) claims a process for the extraction of phosphoric acid from an aqueous solution which comprises treating the solution with an extracting solvent cap able of extracting phosphoric acid from an aqueous solution thereof having a concen-tration of H₁PO, above a threshold value of 35% by weight, whose capacity for ex- must effect the extraction at a relatively low tracting phosphoric acid from water varies temperature and then, taking advantage of inversely with temperature and which does not extract phosphoric acid from an aqueous solution_thereof_having_a_concentration_be 35 low a threshold value of 35% by weight. the temperature of the treatment being low enough for the formation of a clear homogeneous-extract-phase distinct from a residual aqueous-phase; separating the extract phase from the aqueous phase raising the temperature of the extract phase to that at which it separates into a lower layer con-

temperature, and the extracting power of the solvent falls with rising temperature) and (iii) it is not capable of extracting phosphoric acid from an aqueous solution having a concentration below a threshold value

of=35%_by-weight._____ The threshold value is the concentration of acid in water below which the solvent does not preferentially dissolve the acid from the water. The threshold value is not an exact figure, since it can vary with the temperature, and it is different for different solvent, but for the solvents to be used in this process it is about 35% by weight.

To make the process work using a solvent of characteristics (i), (ii) and (iii), one the inverse temperature dependence of the extracting power, separate the extract phase into phosphoric acid and solvent at a tem = perature which is higher than the extracting temperature after removal of the extract phase_from_the_aqueous_solution.

The present invention is an improvement 80 in=or=modification=of=the=process=of=our=said= Application No. 1,112,033, and is concerned with specific solvents useful in carrying out the process of our said earlier application.

PATENT SPECIFICATION

NO DRAWINGS

Inventors .- AVRAGHAM MATITIAJU BANIEL and RUTH BLUMBERG.



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COMPLETE SPECIFICATION

Phosphoric Acid Recovery.

We, Israel Mining Industries Institute for Research and Development, an Israel body corporate of Near Irganim, Haifa—Bay, Israel, do—hereby—declare—the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following ticularly described in and by the following statement:-

phosphoric acid from an aqueous solution containing phosphoric acid. The invention may be used for extracting phosphoric acid from a "technical grade phosphoric acid", 15 i.e. an aqueous phosphoric acid usually This invention relates to the recovery of manufactured by the decomposition of rock
phosphate with sulphuric acid, having an

H₃PO₄ concentration of not less than 35% by weight and possibly up to about 90% 20 by weight, and containing the usual impurities of such phosphoric acid.

Our co-pending Application No. 10235/65 (Serial No. 1,112,033) claims a process for the extraction of phosphoric acid from an 25 aqueous solution which comprises treating the solution with an extracting solvent capable_of_extracting_phosphoric_acid_from=an aqueous solution thereof having a concen-_tration_of=H,PO, above a threshold value 30...of 35% by weight, whose capacity for ex-

tracting phosphoric acid from water varies inversely_with_temperature_and_which_does the temperature of the treatment being low enough for the formation of a clear homo-geneous extract phase distinct from a residual_aqueous_phase, separating the extract

taining phosphoric acid and an upper layer containing solvent, and separating the lower layer from the upper layer. It will be seen, then, that a solvent useful in the process of that application has the following characteristics: (i) it is capable of extracting phosphoric acid from an aqueous solution thereof having a concentration of H,PO, above 50 a "threshold value" of 35% by weight, (ii) its capacity for extracting phosphoric acid from water varies inversely with temperature (that is, the solubility of phosphoric acid in the solvent is greater the lower the temperature of the solvent is greater the lower the temperature, and the extracting power of the solvent falls with rising temperature) and (iii) it is not capable of extracting phosphoric acid from an aqueous solution have ing a concentration below a threshold value of 35% by weight.

The threshold value is the concentration of_acid=in_water_below_which_the_solvent does_not_preferentially_dissolve_the_acid_ from the water. The threshold value is not::65 an exact figure, since it can vary with the temperature, and it is different for different solvent, but for the solvents to be used in this process it is about 35% by weight.

To make the process work using a solvent of characteristics (i), (ii) and (iii), one must effect the extraction at a relatively low temperature and then, taking advantage of the_inverse_temperature_dependence=of_the= extracting power, separate the extract phase into=phosphoric_acid=and=solvent=at=a=tem= perature which is higher than the extracting temperature after removal of the extract phase_from_the_aqueous_solution.

The present invention is an improvement 80 in or modification of the process of our said Application No. 1,112,033, and is concerned phase from the aqueous phase, raising the Application No. 1,112,033, and is concerned temperature of the extract phase to that at with specific solvents useful in carrying out which it separates into a lower layer con. the process of our said earlier application.

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- According to the present invention, therefore, there is provided a process as claimed in-claim=1-of-our-Specification=No.-1,112,033 in which the aqueous solution is extracted sith-cyclohexanone-butyl-acetate, or dibutyl-ether of diethylene glycol, or in which the aqueous solution is extracted with diethylether-and-the-heat-treatment-for-the-separation of the phases is effected under super-

10 atmospheric pressure.
The invention will now be illustrated by the following Examples in which all parts and percentages are by weight:

EXAMPLE 1 (cyclohexanone) 160 g. of wet process phosphoric acid containing 60.5% H₃PO₄ (by wt.) was stirred at 20°C, with 60 g. of cyclohexanone for two liquid phases, which were separated. as well as the greater part of the impurities 25 of the original wet process acid. The solvent phase containing the balance of original phosphoric acid was heated at 65 °C and while being heated was mixed with about 15 ml. of water. Phase separation—

about 15 ml. of water. Phase separation—
about 15 ml. of water. Phase separation—
about 15 ml. of water. The top phase consisted of the solvent. The H₃PO₃ contained—
sisted of the solvent. The H₃PO₃ contained—
in the bottom=was=purified_aqueous=phose—
phoric acid of a concentration of 58%. This purified—phosphoric—acid—was—neutralized—
provided—phosphoric—acid—was—neutralized—
about 15 ml. of water. The top phase concentration of 58%. This purified—phosphoric—acid—was—neutralized—
ammonium—phosphate—without the formaammonium phosphate without the formation-of-any precipitate

butyl acetate for ten minutes, then stirring was stopped. The mixture was allowed to were separated. The upper solvent phase the upper layer.

contained 38 g of H₂PO₄ (calculated as 2. A process as claimed in Claim 1, in 100% H₃PO₄), the balance being in the which the temperature of the extract phase stratify into two liquid phases and the latter lower_phase_together_with_the_impurities_of___is_raised_by_heating_the_extract_to_below_its_110 the original technical grade acid. The solvent extract phase was heated at 45°C. while_being_mixed_with_a small amount of ... water, in order to induce the phase separation. The resultant lower layer, which con-tion of solids during the evaporation.

EXAMPLE 3

containing 51% P.O. were mixed at 25°C with an equal volume of the dibutyl ether of diethylene glycol, resulting in an aqueous phase and 320 g; of solvent extract contain- 65
ing=140 g. H;PO; (calculated as 100%
H;PO;). This extract was decanted and
heated to=100°C; a=small amount of water

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being added to induce phase separation.

The bottom phase comprised 200 g of 70

purified phosphoric acid of 50% P.O. This
acid was concentrated to 95% H.PO. with out-deposition-of-solids-during-evaporation.

> EXAMPLE 4 (diethyl=ether)

50 ml. of wet process phosphoric acid of 70% H-PO concentration were mixed at-25°C. with 100 ml. ethyl ether for 10 minutes::obtaining:115 g. of extract containing 35% H, PO. An aqueous phase containing 80 27% of the original P.O. together with the impurities separated out at the bottom of the_vessel-____

The solvent extract removed was heated to=36°C under pressure to obviate evaporation, thus inducing separation of the extracted P.O. as a clean phosphoric acid.

WHAT WE CLAIM IS: 1. A process for the extraction of phosphoric acid from an aqueous solution containing phosphoric acid, which comprises treating the aqueous solution with cyclohexanone, butyl acetate, dibutyl ether of diethylene glycol, or diethyl ether, the temperature of the treatment being low enough 95 for the formation of a clear homogeneous extract phase distinct from a residual aque-ous phase, separating the organic extract phase from the aqueous phase, raising the temperature of the extract phase to that at 100 which it separates into a lower layer containing phosphoric acid and an upper layer containing solvent, said raising of the temperature being effected under superatmos pheric pressure when the solvent is diethyl 105 ether, and separating the lower layer from-

3....A process for the extraction of phosphoric: acid from an aqueous solution thereof, substantially as described in Example 1,

3 or 4:

4. Phosphoric acid obtained by a pro-

J.A. KEMP & CO., EXAMPLE 3 Cliartered Patent Agents,

idibutylether of diethylene glycol) = 14. South Square. Gray's Inn

200 ml. of wet process phosphoric acid London, W.C.I.

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